L1	STRUCTURE UPLOADED
L2	0 S L1
L3	3 S L1 SSS FULL
	FILE 'HCAPLUS' ENTERED AT 10:36:13 ON 16 OCT 2008
L4	22 S L3
L5	11 S L4 AND (PY<2003 OR AY<2003 OR PRY<2003)
	FILE 'HCAPLUS' ENTERED AT 13:09:49 ON 16 OCT 2008
L1	FILE 'HCAPLUS' ENTERED AT 13:09:49 ON 16 OCT 2008 77099 S (TMF(W) (ALPHA OR .ALPHA)) OR ((TUMOR NECROSIS FACTOR)(W) (ALPH
L1 L2	
	77099 S (TNF(W)(ALPHA OR .ALPHA)) OR ((TUMOR NECROSIS FACTOR)(W)(ALPH
L2	77099 S (TNF(W)(ALPHA OR .ALPHA)) OR ((TUMOR NECROSIS FACTOR)(W)(ALPH 77099 S (TNF(W)(ALPHA OR A)) OR ((TUMOR NECROSIS FACTOR)(W)(ALP

FILE 'REGISTRY' ENTERED AT 10:35:37 ON 16 OCT 2008

=> file registry COST IN U.S. DOLLARS

SINCE FILE TOTAL. ENTRY SESSION 0.21 0.21

FULL ESTIMATED COST

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REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/support/stngen/stndoc/properties.html

Uploading C:\Program Files\STNEXP\Queries\10693722specific.str

chain nodes :

10 11 12 13 20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37

ring nodes :

1 2 3 4 5 6 7 8 9 14 15 16 17 18 19

chain bonds :

 $1-22 \quad 2-34 \quad 3-33 \quad 4-32 \quad 7-26 \quad 8-10 \quad 9-25 \quad 10-11 \quad 10-14 \quad 10-28 \quad 11-12 \quad 11-36 \quad 11-37 \quad 11-37 \quad 11-38 \quad 11-$ 12-13 15-31 16-21 17-20 18-30 19-29 22-23 22-35 23-24 23-27 ring bonds :

 $1-2 \quad 1-6 \quad 2-3 \quad 3-4 \quad 4-5 \quad 5-6 \quad 5-7 \quad 6-9 \quad 7-8 \quad 8-9 \quad 14-15 \quad 14-19 \quad 15-16 \quad 16-17 \quad 17-18$ 18-19 exact/norm bonds : 1-22 5-7 6-9 7-8 7-26 8-9 8-10 9-25 22-23 23-24

exact bonds :

2-34 3-33 4-32 10-11 10-14 10-28 11-12 11-36 11-37 12-13 15-31 16-21

17-20 18-30 19-29 22-35 23-27 normalized bonds :

1-2 1-6 2-3 3-4 4-5 5-6 14-15 14-19 15-16 16-17 17-18 18-19

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:CLASS 11:CLASS 12:CLASS 13:CLASS 14:Atom 15:Atom 16:Atom 17:Atom 18:Atom 19:Atom

20:CLASS 21:CLASS

22:CLASS 23:CLASS 24:CLASS 25:CLASS 26:CLASS 27:CLASS 28:CLASS 29:CLASS 30:CLASS 31:CLASS

32:CLASS 33:CLASS 34:CLASS 35:CLASS 36:CLASS 37:CLASS

## L1 STRUCTURE UPLOADED

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SAMPLE SEARCH INITIATED 10:35:59 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 5 TO ITERATE

100.0% PROCESSED 5 ITERATIONS

0 ANSWERS

SEARCH TIME: 00.00.01

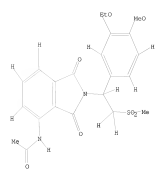
FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\* BATCH \*\*COMPLETE\*\* PROJECTED ITERATIONS: 5 TO 234 PROJECTED ANSWERS: 0 TO 0

L2 0 SEA SSS SAM L1

=> d 11

L1 HAS NO ANSWERS

STR



Structure attributes must be viewed using STN Express query preparation.

3 ANSWERS

=> s 11 sss full FULL SEARCH INITIATED 10:36:08 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 122 TO ITERATE

100.0% PROCESSED 122 ITERATIONS SEARCH TIME: 00.00.01

L3 3 SEA SSS FUL L1

=> file hcaplus

 COST IN U.S. DOLLARS
 SINCE FILE
 TOTAL

 ENTRY
 SESSION

 FULL ESTIMATED COST
 178.36
 178.57

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FILE COVERS 1907 - 16 Oct 2008 VOL 149 ISS 16 FILE LAST UPDATED: 15 Oct 2008 (20081015/ED) HCAplus now includes complete International Patent Classification (IPC) reclassification data for the second quarter of 2008.

New CAS Information Use Policies, enter HELP USAGETERMS for details.

This file contains CAS Registry Numbers for easy and accurate substance identification.

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=> s 13

L4 22 L3

=> s 14 and (PY<2003 or AY<2003 or PRY<2003)

22959050 PY<2003

4498362 AY<2003

3966940 PRY<2003

L5 11 L4 AND (PY<2003 OR AY<2003 OR PRY<2003)
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 $\Rightarrow$  d 15 1-11 ti abs bib

L5 ANSWER 1 OF 11 HCAPLUS COPYRIGHT 2008 ACS on STN

TI Solid forms of (+)-2-[1-(3-ethoxy-4-methoxyphenyl)-2-methylsulfonylethyl)-4-acetylaminoisoindoline-1,3-dione, compositions thereof, and uses thereof AB Solid forms comprising (+)-2-[1-(3-Ethoxy-4-methoxyphenyl)-2-

methylsulfonylethyll-4-acetylaminoisoindoline-1,3-dione (1), compns. comprising the solid forms, methods of making the solid forms and methods of their use are disclosed. The methods include methods of treating and/or preventing disorders ameliorated by the reduction of levels of TNF-a or the inhibition of PDE4. I was prepared by the reaction of 1-(3-ethoxy-4-methoxyphenyl)-2-methylsulfonylethylamine with 3-acetamidophthalic anhydride, yield = 59%.

AN 2008:1156159 HCAPLUS <<LOGINID::20081016>>

To Solid forms of (+)-2-[1-(3-ethoxy-4-methoxypheny1)-2-methylsulfonylethyl]-4-acetylaminoisoindoline-1,3-dione, compositions thereof, and uses thereof IN Muller, George W; Schafer, Peter H; Man, Hon-Wah; Ge, Chuansheng; Xu,

Jean PA USA

SO U.S. Pat. Appl. Publ., 66pp., Cont.-in-part of U.S. Ser. No. 106,142.

CODEN: USXXCO DT Patent

LA English

FAN.CNT 4

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	US 20080234359	A1	20080925	US 2008-79615	20080327 <
	US 20030187052	A1	20031002	US 2003-392195	20030319 <
	US 6962940	B2	20051108		
	CN 1965823	A	20070523	CN 2006-10137407	20030320 <
	US 20050192336	A1	20050901	US 2005-106142	20050413 <
	US 7427638	B2	20080923		
	US 20050267196	A1	20051201	US 2005-170308	20050628 <
	US 7358272	B2	20080415		
	US 20080027123	A1	20080131	US 2007-824523	20070629 <
	US 20080207730	A1	20080828	US 2008-69282	20080208 <
	US 20080242719	A1	20081002	US 2008-98379	20080404 <
PRAI	US 2002-366515P	P	20020320	<	
	US 2003-438450P	P	20030107		
	US 2003-392195	A3	20030319		
	US 2005-106142	A2	20050413		
	CN 2003-811093	A3	20030320		
	US 2005-170308	A3	20050628		

- L.5 ANSWER 2 OF 11 HCAPLUS COPYRIGHT 2008 ACS on STN
- Compositions comprising selective cytokine inhibitory drugs for treatment and management of macular degeneration and Methods of using thereof
- AB Methods of treating, preventing and/or managing macular degeneration are disclosed. Specific embodiments encompass the administration of a selective cytokine inhibitory drug, or a pharmaceutically acceptable salt, solvate, hydrate, stereoisomer, clathrate, or prodrug thereof, alone or in combination with a second active agent and/or surgery. Pharmaceutical compns., single unit dosage forms, and kits suitable for use in methods of the invention are also disclosed. Thus, patients with macular degeneration received conventional therapy with verteporfin and (+)-2-[1-(3-ethoxy-4 methoxyphenyl)-2-methylsulfonylethyl]-4 acetylaminoisoindoline 1,3-dione in an amount of about 20 mg/day as an adjuvant for 20 wk. The neovascular cascade was sufficiently hindered in those patients to indefinitely prolong the effects of the photodynamic therapy.
- 2007:998162 HCAPLUS <<LOGINID::20081016>> AN
- DN 147:330440
- ΤI Compositions comprising selective cytokine inhibitory drugs for treatment and management of macular degeneration and Methods of using thereof
- IN Zeldis, Jerome B. PA USA
- SO U.S. Pat. Appl. Publ., 30pp., Cont.-in-part of U.S. Ser. No. 699,110.
- CODEN: USXXCO DT Patent
- T.A English
- FAN.CNT 2

		PATENT NO.				KIND DATE									DATE					
P	I	US 20070207121 US 20040091454 WO 2005044269 W: AE, AG, AL				A1 20040513 A1 20050519				US 2	006- 003-	5761 6991	40 10		2		030 <-			
				CN, GE, LK, NO, TJ, BW, AZ, EE, SI,	CO, GH, LR, NZ, TM, GH, BY, ES,	CR, GM, LS, OM, TN, GM, KG, FI, TR,	CU, HR, LT, PG, TR, KE, KZ, FR,	CZ, HU, LU, PH, TT, LS, MD, GB,	DE, ID, LV, PL, TZ, MW, RU, GR,	DK, IL, MA, PT, UA,	DM, IN, MD, RO, UG, NA, TM, IE,	DZ, IS, MG, RU, US, SD, AT, IT,	EC, JP, MK, SC, UZ, SL, BE, LU,	EE, KE, MN, SD, VC, SZ, BG, MC,	EG, KG, MW, SE, VN, TZ, CH, NL,	ES, KP, MX, SG, YU, UG, CY, PL,	FI, KR, MZ, SK, ZA, ZM, CZ, PT,	GB, KZ, NA, SL, ZM, ZW, DE, RO,	GD, LC, NI, SY, ZW AM, DK, SE,	
		US WO US AU	2008 2003 2004 2002 2003	2014 -699 -US1 -422 -285	18 110 3253 900P 107		W P		2003 2004	0428 1031			008-	2014	18		2	0080	327	
0	5	MAI	RPAT	T4/:	3304	40														

- L5 ANSWER 3 OF 11 HCAPLUS COPYRIGHT 2008 ACS on STN
- Methods of the treatment or prevention of exercise-induced asthma using (+)-2-[1-(3-ethoxy-4-methoxyphenyl)-2-methylsulfonylethyl]-4acetylaminoisoindoline-1.3-dione
- Methods of treating, managing or preventing exercise-induced asthma are disclosed. Specific methods encompass the administration of (+)-2-[1-(3-ethoxy-4-methoxypheny1)-2-methylsulfonylethyl]-4acetylaminoisoindoline-1,3-dione alone or in combination with a second active agent. Pharmaceutical compns. and single unit dosage forms are also disclosed.

- 2006:823362 HCAPLUS <<LOGINID::20081016>> AN
- DN 145:224862
- ΤТ Methods of the treatment or prevention of exercise-induced asthma using (+)-2-[1-(3-ethoxy-4-methoxyphenyl)-2-methylsulfonylethyl]-4acetylaminoisoindoline-1,3-dione
- Muller, George W.; Schafer, Peter H.; Rohane, Patricia E. W. IN
- Celgene Corporation, USA PA
- SO U.S. Pat. Appl. Publ., 32pp., Cont.-in-part of U.S. Ser. No. 106,142.
- CODEN: USXXCO DT Patent
- T.A
- English FAN.CNT 4

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	US 20060183788	A1	20060817	US 2006-392846	20060328 <
	US 7276529	B2	20071002		
	US 20030187052	A1	20031002	US 2003-392195	20030319 <
	US 6962940	B2	20051108		
	CN 1965823	A	20070523	CN 2006-10137407	20030320 <
	US 20050192336	A1	20050901	US 2005-106142	20050413 <
	US 7427638	B2	20080923		
	US 20050267196	A1	20051201	US 2005-170308	20050628 <
	US 7358272	B2	20080415		
	US 20080027123	A1	20080131	US 2007-824523	20070629 <
	US 20080207730	A1	20080828	US 2008-69282	20080208 <
	US 20080242719	A1	20081002	US 2008-98379	20080404 <
PRAI	US 2002-366515P	P	20020320	<	
	US 2003-438450P	P	20030107		
	US 2003-392195	A3	20030319		
	US 2005-106142	A2	20050413		
	CN 2003-811093	A3	20030320		
	US 2005-170308	A.3	20050628		

- RE.CNT 131 THERE ARE 131 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- T. 5 ANSWER 4 OF 11 HCAPLUS COPYRIGHT 2008 ACS on STN
- TΙ Methods of the treatment of psoriatic arthritis using
  - (+) -2-[1-(3-ethoxy-4-methoxyphenyl)-2-methylsulfonylethyl]-4acetylaminoisoindoline-1,3-dione
- AB Methods of treating, managing or preventing psoriatic arthritis are disclosed. Specific methods encompass the administration of (+)-2-[1-(3-ethoxy-4-methoxyphenyl)-2-methylsulfonylethyl]-4acetylaminoisoindoline-1.3-dione alone or in combination with a second active agent. Pharmaceutical compns. and single unit dosage forms are
- also disclosed. 2006:821184 HCAPLUS <<LOGINID::20081016>> AN
- DN 145:224861
- TI Methods of the treatment of psoriatic arthritis using (+)-2-[1-(3-ethoxy-4-methoxyphenyl)-2-methylsulfonylethyl]-4acetylaminoisoindoline-1,3-dione
- IN Muller, George W.; Schafer, Peter H.; Rohane, Patricia E. W.
- PA Celgene Corporation, USA
- U.S. Pat. Appl. Publ., 19pp., Cont.-in-part of U.S. Ser. No. 106,142. SO CODEN: USXXCO
- Patent
- LA English
- DAM ONT 4

L 11114.	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ΡI	US 20060183787	A1	20060817	US 2006-392845	20060328 <
	US 7208516	B2	20070424		

	US	20030187052	A1	20031002	US	2003-392	195	20030319	<
	US	6962940	B2	20051108					
	CN	1965823	A	20070523	CN	2006-101	37407	20030320	<
	US	20050192336	A1	20050901	US	2005-106	142	20050413	<
	US	7427638	B2	20080923					
	US	20050267196	A1	20051201	US	2005-170	308	20050628	<
	US	7358272	B2	20080415					
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	US	20080207730	A1	20080828	US	2008-692	82	20080208	
	US	20080242719	A1	20081002	US	2008-983	79	20080404	<
PRAI		2002-366515P	P	20020320	<				
	US	2003-438450P	P	20030107					
	US	2003-392195	A3	20030319					
		2005-106142	A2	20050413					
	CN	2003-811093	A3	20030320					
		2005-170308	A3	20050628					
RE.CI	T	130 THERE AR	E 130 CI	TED REFEREN	CES AV	VAILABLE	FOR THIS	RECORD	

ALL CITATIONS AVAILABLE IN THE RE FORMAT

- ANSWER 5 OF 11 HCAPLUS COPYRIGHT 2008 ACS on STN
- ΤI Preparation of substituted phenethyl sulfones and methods of reducing TNFα levels

GI

- The title compds. I [Y = CO, CH2, SO2, CH2C(O); R1-R4 = H, halo, alkyl, alkoxy, etc.; R5, R6 = H, alkyl, alkoxy, CN, etc.; R7 = OH, alkyl, Ph, etc.], useful for reducing TNFa levels and treating inflammatory and autoimmune diseases, were prepared and formulated. E.g., a 2-step synthesis of 2-[1-(3-ethoxy-4-methoxypheny1)-2-methylsulfonylethyl]isoindolin-1-one, starting from di-Me sulfone and 3-ethoxy-4-methoxybenzaldehyde, was given.
- ΑN 2006:425851 HCAPLUS <<LOGINID::20081016>> DN 147:189068
- Preparation of substituted phenethyl sulfones and methods of reducing TNFa levels
- IN Man, Hon-Wah; Muller, George W.
- PA Celgene Corporation, USA
- SO Aust. Pat. Appl., 53 pp.
- CODEN: AUXXCM
- Patent

ΤI

LA English FAN CNT 3

E Pars.	CNI				
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PΙ	AU 2006200033	A1	20060202	AU 2006-200033	20060106
	AU 2006200033	B2	20080814		
	AU 2003203681	A1	20030703	AU 2003-203681	20030409 <

	AU 2003203681	B2	20051006	
PRAI	AU 2003-203681	A3	20030409	
	AU 2000-14472	A3	19991019	<
	WO 1999-US24376	W	19991019	<
os	CASREACT 147:189068			

- ANSWER 6 OF 11 HCAPLUS COPYRIGHT 2008 ACS on STN L5
- Methods of using and compositions comprising selective cytokine inhibitory drugs for treatment and management of macular degeneration
- Methods of treating, preventing and/or managing macular degeneration are disclosed. Specific embodiments encompass the administration of a selective cytokine inhibitory drug, or a pharmaceutically acceptable salt, solvate, hydrate, stereoisomer, clathrate, or prodrug thereof, alone or in combination with a second active agent and/or surgery. Pharmaceutical compns., single unit dosage forms, and kits suitable for use in methods of the invention are also disclosed. Patients with macular degeneration were treated by photodynamic therapy with verteporfin alone, or with the addition of 20 mg/day of selective cytokine inhibitory drug (+)-2-[1-(3-ethoxy-4 methoxyphenyl)-2-methylsulfonylethyl]-4 acetylaminoisoindoline 1,3-dione. The neovascular cascade is sufficiently hindered in the group receiving (+)-2-[1-(3-ethoxy-4 methoxypheny1)-2-methylsulfonylethyl]-4 acetylaminoisoindoline 1,3-dione to indefinitely prolong the effects of the photodynamic therapy.
- 2004:392056 HCAPLUS <<LOGINID::20081016>> AN
- DN 140:386062
- Methods of using and compositions comprising selective cytokine inhibitory drugs for treatment and management of macular degeneration
- IN Zeldis, Jerome B.
- PA
- SO U.S. Pat. Appl. Publ., 19 pp.
- CODEN: USXXCO Patent
- DT LA. English

FAN.CNT 2 PATENT NO. KIND DATE APPLICATION NO. DATE																		
PA:	ENT	NO.			KIN	D	DATE					ION :				ATE		
US CA WO	2004 2504 2004 2004	0091 263 0411	454 81		A1 A1 A2		2004	0521 0521		CA 2	003-	2504	263		2	00310 00310 00310	031	<
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BR	2003	0158	89		A		2005	1004		BR 2	003-	1588	9		2	0031	J31	<
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JΡ	2006509743				T 20060323			8 CN 2003-80108090 3 JP 2004-550274						2	0031	331	<	
									1 NZ 2003-540185									
	AU 2004286824 CA 2543618						9 AU 2004-286824											
CA	2543	618						9 CA 2004-2543618					20040428					

	WO	2005	0442	69	A1 20050519 AL, AM, AT, AU, AZ,					9 WO 2004-US13253										
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												SC,								
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				TD,																
	EP	1684								EP 2004-750923						20040428 , SE, MC, PT,				
		R:																		
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		2004		70		A			0123			004-								
		1901				A			0124			004-					0040			
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		2007				A1			0906			006-								
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		2003				W		2003												
		2004				W		2004	0428											
OS	MAI	RPAT	140:	3860	52															

L5 ANSWER 7 OF 11 HCAPLUS COPYRIGHT 2008 ACS on STN

TI Use of (+)-2-[1-(3-ethoxy-4-methoxypheny1)-2-methylsulfonylethyl]-4-acetylaminoisoindoline-1,3-dione and compositions thereof for inhibiting TNF- $\alpha$  production and PDE4 activity

AB The invention discloses stereomerically pure
(S)-2-[1-(3-Ethoxy-4-methoxyphenyl)-2-methylsulfonylethyl]-4acetylaminoisoindoline-1,3-dione (+)-I, substantially free of its
(-)-isomer, and prodrugs, metabolites, polymorphs, salts, solvates,
hydrates, and clathrates thereof. Methods of using and pharmaceutical
compns. comprising (+)-I for treating and/or preventing disorders
ameliorated by the reduction of levels of tumor necrosis factor α
(TNF-α) or the inhibition of phosphodiesterase IV (PDE4) are also
disclosed. Examples include the synthesis and resolution of (+)-I, thirteen

bioassays, an aqueous solubility study, and three formulations. For instance, 3-nitrophthalic acid was hydrogenated using 10% Pd/C in BtOH to give the amine (84%), which was condensed with Ac2O to afford 3-acetamidophthalic anhydride (61%). Reaction of the phthalic anhydride with 1-(3-ethoxy-4-methoxyphenyl)-2-(methylsulfonyl)ethylamine to give I (59%), followed by resolution with N-acetyl-L-leucine in MeOH provided (+)-I (90% recovery, 98.4% ee). The latter inhibited LPS-induced TNP-a production by human whole blood and PDE4 activity with IC5O values of 294 nM and 73.5 nM, resp. (+)-I showed >500-fold to >40,000-fold selectivity for PDE4 over PDE1, PDE2, PDE3, PDE3, and PDE6. In addition, (+)-I suppressed LPS-induced lung neutrophilia in conscious ferrets with an ED50 of 0.8 mg/kg. Thus, (+)-I and its pharmaceutical compns. are useful for treating and/or preventing cancer, depression, and a variety of allergic, inflammatory, and autoimmune disorders (no data).

- AN 2003:777583 HCAPLUS <<LOGINID::20081016>>
- DN 139:296870
- TI Use of (+)-2-[1-(3-ethoxy-4-methoxyphenyl)-2-methylsulfonylethyl]-4acetylaminoisoindoiine-1, 3-dione and compositions thereof for inhibiting TNF-a production and PDE4 activity
- IN Schafer, Peter H.; Muller, George W.; Man, Hon-Wah; Ge, Chuansheng PA Celgene Corporation, USA
- PA Celgene Corporation, U SO PCT Int. Appl., 57 pp.
- CODEN: PIXXD2
- DT Patent
- LA English
- FAN CNT 4

PATENT NO.							TND DATE				ADDITORTION NO					D3.000				
	PATENT NO.																ATE			
PΙ	WO																	320 <		
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		P	L,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	ТJ,	TM,	TN,	TR,	TT,	TZ,		
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	ΑU	200322	29		A1		2003	1008		AU 2	003-	2247		20030320 <						
	ΑU	200322	2003224729				B2 20080103													
	EΡ	148508								EP 2003-721414										
																		PT,		
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	CN	165277	2			A		2005	0810		CN 2	003-	8110	93		2	0030	320 < 320 <		
	JP	200552	538	36		T		2005	0825		JP 2	003-	5778	77		2	0030	320 <		
	NZ	535798				A		2006	0428		NZ 2	003-	5357	98		2	0030:	320 < 320 <		
	CN	196582	:3			A		2007	0523		CN 2	006-	1013	7407		2	0030	320 <		
		2004PA				A		2005	0713		MX 2	004-	PA90	75		2	00409	920 <		
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	US	200802	07	730		A1		2008	0828									208 <		
		200802						2008	1002		US 2	008-	9837	9		2	0800	404 <		
PRAI	US	JS 2002-366515P				P		2002	0320	<-	-									
	US	2003-4	384	450P		P		2003	0107											
	US	US 2003-392195				A3		2003	0319											
	CN 2003-811093						2003	0320												
					W		2003	0320												
	WO 2003-US8738 US 2005-170308					A3		2005	0628											

RE.CNT 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L5 ANSWER 8 OF 11 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI Use of (-)-2-[1-(3-ethoxy-4-methoxyphenyl)-2-methylsulfonylethyl]-4-acetylaminoisoindoline-1,3-dione and compositions thereof for inhibiting TNF- $\alpha$  production and PDB4 activity

GT

AB The invention discloses stereomerically pure (R)-2-[1-(3-Ethoxy-4-methoxyphenyl)-2-methylsulfonylethyl]-4acetylaminoisoindoline-1,3-dione (-)-I, substantially free of its (+)-isomer, and prodrugs, metabolites, polymorphs, salts, solvates, hydrates, and clathrates thereof. Methods of using and pharmaceutical compns. comprising (-)-I for treating and/or preventing disorders ameliorated by the reduction of levels of tumor necrosis factor  $\alpha$  $(TNF-\alpha)$  or the inhibition of phosphodiesterase IV (PDE4) are also disclosed. Examples include the synthesis and resolution of (-)-I, seven bioassays, an aqueous solubility study, and three formulations. For instance, 3-nitrophthalic acid was hydrogenated using 10% Pd/C in EtOH to give the amine (84%), which was condensed with Ac20 to afford 3-acetamidophthalic anhydride (61%). Reaction of the phthalic anhydride with 1-(3-ethoxy-4-methoxyphenyl)-2-(methylsulfonyl)ethylamine to give I (59%), followed by resolution with N-acetyl-D-leucine in MeOH provided (-)-I (90% recovery, 98.4% ee). The latter inhibited LPS-induced TNF-a production by human whole blood and PDE4 activity with IC50 values of 371 nM and 611 nM, resp. (-)-I showed >45-fold to >39,000-fold selectivity for PDE4 over PDE1, PDE2, PDE3, PDE5, and PDE6. Thus, (-)-I and its pharmaceutical compns. are useful for treating and/or preventing cancer, depression, and

AN 2003:777582 HCAPLUS <<LOGINID::20081016>>

- DN 139:296869
- II Use of (-)-2-[1-(3-ethoxy-4-methoxyphenyl)-2-methylsulfonylethyl]-4-acetylaminoisoindoline-1,3-dione and compositions thereof for inhibiting TNF- $\alpha$  production and PDB4 activity

a variety of allergic, inflammatory, and autoimmune disorders (no data).

- IN Schafer, Peter H.; Muller, George W.; Man, Hon-Wah; Ge, Chuansheng
- PA Celgene Corporation, USA
- SO PCT Int. Appl., 49 pp.
- CODEN: PIXXD2
- DT Patent
- LA English
- FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

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A1 20031002 W0 2003-US8737 20030320 <--
PΤ
   WO 2003080048
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
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            PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ,
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            BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
    AU 2003222034 A1 20031008 AU 2003-222034
                                                               20030320 <--
PRAI US 2002-366516P
                       P
                             20020320 <--
    US 2003-438448P
WO 2003-US8737
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                             20030320
RE.CNT 3
            THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD
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- ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L5 ANSWER 9 OF 11 HCAPLUS COPYRIGHT 2008 ACS on STN TI Interactions between myeloma and endothelial cells and the effects of thalidomide and its analogues
- AB Modeling the situation observed in vivo, the authors examined the effect of thalidomide and its analogs in co-cultures of myeloma and endothelial cells. It was found that myeloma cells in co-culture had significantly lower levels of CC-1004- and CC-1088-induced apoptosis than those cultured alone. Interestingly, basal apoptosis was also lower in RPMI-8226/S co-cultured with endothelial cells compared to myeloma cell culture. The authors' data suggest that myeloma/endothelial cell interactions in co-culture have a significant protective effect on both basal and drug-induced levels of apoptosis in myeloma cells.
- AN 2003:649755 HCAPLUS <<LOGINID::20081016>>
- DN 140:228565
- TI Interactions between myeloma and endothelial cells and the effects of thalidomide and its analogues
- AU Molostvov, G.; Morris, A.; Rose, P.; Basu, S.
- CS University of Warwick, Coventry, UK
- SO Free Papers Annual Meeting of the European Haematology Association, 7th, Florence, Italy, June 6-9, 2002 (2002), 263-266 Publisher: Monduzzi Editore, Bologna, Italy. CODEN: 69EIOR; ISBN: 88-323-2606-X
- DT Conference
- LA English
- RE.CNT 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD
  ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L5 ANSWER 10 OF 11 HCAPLUS COPYRIGHT 2008 ACS on STN
- TI Preparation of substituted phenethylsulfones for reducing  $\text{TNF}\alpha$  levels

AB The title compds. [I] the carbon atom designated "\*" constitutes a center of chirality; Y = CO, CH2 CH2CO; R1-R4 = H, halo, alkyl, etc.; R5, R6 = H, alkyl, alkoxy, etc.; R7 = OH, alkyl, Ph, etc.] which reduce the levels of TNFw and inhibit PDE IV in a mammal (no data), were prepared and formulated. Typical embodiments are 2-[1-(3-ethoxy-4-methoxyphenyl)-2-methylsulfonylethyl]-4-aminoisoindoline-1,3-dione and 2-[1-(3-eytlopentyloxy-4-methoxyphenyl)-2-

methylsulfonylethyl]isoindoline-1,3-dione. 2000:78904 HCAPLUS <<LOGINID::20081016>>

AN 2000:78904 DN 132:107873

TI Preparation of substituted phenethylsulfones for reducing TNF $\alpha$  levels

Ι

IN Muller, George W.; Man, Hon-wah

PA Celgene Corporation, USA

SO U.S., 13 pp.

CODEN: USXXAM

DT Patent

LA English

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	JP	20025	52849	96		T		2002	0903		JP 2	000-	5792	18		1	9991	019	<
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	NO 319790					B1		2005											
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PRAI	PRAI US 1998-183049					A3		1998											

EP 1999-971317 A.3 19991019 <--WO 1999-US24376 19991019 <--

MARPAT 132:107873 OS

THERE ARE 29 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 29 ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 11 OF 11 HCAPLUS COPYRIGHT 2008 ACS on STN 1.5

Ι

ΤI Preparation of substituted phenethylsulfones and method of reducing TNFa levels

- The title compds. [I; the carbon atom designated \* constitutes a center of AB chirality; Y = SO2, CO, CH2; R1-R4 = H, halo, alkyl, etc.; R5, R6 = H, alkyl, alkoxy, etc.; R7 = OH, alkyl, Ph, etc.], useful in reducing the levels of  $TNF\alpha$  and inhibiting PDE IV (no data), were prepared and formulated. Typical embodiments are 2-[1-(3-ethoxy-4-methoxypheny1)-2-methylsulfonylethyl]-4-aminoisoindoline-
  - 1,3-dione and 2-[1-(3-cyclopentyloxy-4-methoxyphenyl)-2-
- methylsulfonylethyl]isoindoline-1,3-dione (prepns. were given). 2000:10631 HCAPLUS <<LOGINID::20081016>> AN
- DM 132:64167
- ΤI Preparation of substituted phenethylsulfones and method of reducing TNFa levels
- IN Muller, George W.; Man, Hon-Wah
- PA Celgene Corporation, USA
- SO U.S., 12 pp., Division of U.S. Ser. No. 183,049.
- CODEN: USXXAM DT Patent
- LA English

FAN.CNT 3				
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI US 6011050	A	20000104	US 1999-340617	19990629 <
US 6020358	A	20000201	US 1998-183049	19981030 <
PRAI US 1998-183049	A3	19981030	<	
OS MARPAT 132:64167				
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RE.CNT 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

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             2 SYSTROPHY
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         74307 REGIONAL
         63422 PAIN
        149117 SYNDROME
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=> s 14 and (PY<2003 or AY<2003 or PRY<2003)
      22959050 PY<2003
       4498362 AY<2003
       3966940 PRY<2003
             4 L4 AND (PY<2003 OR AY<2003 OR PRY<2003)
=> d 15 1-4 ti abs bib
     ANSWER 1 OF 4 HCAPLUS COPYRIGHT 2008 ACS on STN
     Methods of using and compositions comprising selective cytokine inhibitory
     drug for treatment, modification and management of pain
     Methods of treating, preventing, modifying and managing various types of
AB
     pain are disclosed. Specific methods comprise the administration of a
     selective cytokine inhibitory drug, or a pharmaceutically acceptable salt,
     solvate, hydrate, stereoisomer, clathrate, or prodrug thereof, alone or in
     combination with a second active agent and/or surgery, psychol. or phys.
     therapy. Pharmaceutical compns., single unit dosage forms, and kits
     suitable for use in methods of the invention are also disclosed.
AN
    2005:426388 HCAPLUS <<LOGINID::20081016>>
DN
TΙ
     Methods of using and compositions comprising selective cytokine inhibitory
     drug for treatment, modification and management of pain
     Zeldis, Jerome B.; Faleck, Herbert; Manning, Donald C.
TN
PA
    Celgene Corporation, USA
    PCT Int. Appl., 85 pp.
SO
     CODEN: PIXXD2
DT
     Patent
LA
    English
FAN.CNT 6
     PATENT NO.
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                                                                   DATE
     WO 2005043971
                         A2
                                20050519
                                           WO 2004-US12722
                                                                   20040423
                               20050714
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             CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
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                                             MX 2006-PA4381
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     US 20070161696
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                                             US 2007-576139
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PRAI US 2003-693794
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     US 2003-693722
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     WO 2004-US12722
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     MARPAT 142:457121
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L5 ANSWER 2 OF 4 HCAPLUS COPYRIGHT 2008 ACS on STN
TI Preparation of 2-(fluoroalkoxyphenylalkyl)-1.3-di

TI Preparation of 2-(fluoroalkoxyphenylalkyl)-1,3-dihydroisoindolones as PDE4, TNF- $\alpha$  , and/or MMP inhibitors

$$X^4$$
  $O$   $O-R^2$   $X^3$   $Y$   $Y$   $Z$ 

AB Title compds. I [wherein X1-X4 = independently H, halo, NO2, NH2, CF3, alkyl, cycloalkyl(alkyl), NR7R8-(alkyl), R8CONH-(alkyl), NR7R8CONH-(alkyl), R8CONH-(alkyl), midiazolyl(alkyl), pyrrolyl(alkyl), oxadiazolyl(alkyl), triazolyl(alkyl); or X1 and X2 or X2 and X3 or X3 and X4 may be taken together to form a (hetero)cycloalkyl ring; Y = CO, CH2, CH2CO, COCH2, SO2; Z = H, COR3, alkylsulfonyl(alkyl),

ΙI

alkyl, CH2OH, alkoxymethyl, CN; R1 and R2 = independently CHF2, alkyl, cycloalkyl(alkyl); at least one of R1 and R2 = CHF2; R3 = NR4R5, alkyl, OH, alkoxy, (un) substituted Ph, PhCH2; R4 and R5 = independently H, alkyl, OH, OCOR6; R6 = alkyl(amino), Ph, PhCH2, aryl; R7 and R8 = independently H, alkyl, cycloalkyl(alkyl), NR7R8-alkyl, R80-alkyl, Ph, PhCH2, aryl; or pharmaceutically acceptable salts, hydrates, solvates, clathrates, stereoisomers, and prodrugs thereof] were prepared For example, alkylation of 3,4-dihydroxybenzaldehyde with chlorodifluoromethane in the presence of K2CO3 in DMF gave 4-difluoromethoxy-3-hydroxybenzaldehyde (15%), which was further alkylated with bromomethylcyclopropane under the same conditions to afford 3-cyclopropylmethoxy-4-difluoromethoxybenzaldehyde (100%). Reaction of the benzaldehyde with ammonium acetate in 95% EtOH, followed by addition of malonic acid provided 3-amino-3-(3-cyclopropylmethoxy-4difluoromethoxyphenyl)propionic acid (52%). Condensation of the amine with 3-acetamidophthalic anhydride using sodium acetate in AcOH yielded the isoindoledione II (85%). I and their pharmaceutical compns., optionally in combination with another therapeutic agent, are useful for the treatment or prevention of diseases associated with phosphodiesterase 4 (PDE4) inhibition, abnormal tumor necrosis factor  $\alpha$  (TNF- $\alpha$  ) levels

, and/or matrix metalloproteinase (MMP) inhibition, such as myelodysplastic syndrome, myeloproliferative disease, complex regional pain syndrome, cancer, inflammatory

diseases, and autoimmune diseases (no data).
AN 2004:589381 HCAPLUS <<LOGINID::20081016>>

DN 141:140314

TI Preparation of 2-(fluoroalkoxyphenylalkyl)-1,3-dihydroisoindolones as PDE4, TNF- $\alpha$ , and/or MMP inhibitors

IN Muller, George W.; Man, Hon-Wah; Zhang, Weihong

PA Celgene Corporation, USA

SO PCT Int. Appl., 98 pp.

CODEN: PIXXD2

DT Patent LA English FAN.CNT 1

	PATENT NO.					KIND DATE			APPLICATION NO.											
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WO 2003-US41568 W 20031229 MARPAT 141:140314 OS ANSWER 3 OF 4 HCAPLUS COPYRIGHT 2008 ACS on STN L5 TI Evidence for local inflammation in complex regional pain syndrome type 1 AB BACKGROUND: The pathophysiol, of complex regional pain syndrome type 1 (CRPS 1) is still a matter of debate. Peripheral afferent, efferent and central mechanisms are supposed. Based on clin. signs and symptoms (e.g. edema, local temperature changes and chronic pain) local inflammation is suspected. Aim: To determine the involvement of neuropetides, cytokines and eicosanoids as locally formed mediators of inflammation. Methods: In this study, nine patients with proven CRPS 1 were included. Disease activity and impairment was determined by means of a Visual Analog Scale, the McGill Pain Questionnaire, the difference in volume and temperature between involved and uninvolved extremities, and the reduction in active range of motion of the involved extremity. Venous blood was sampled from and suction blisters made on the involved and uninvolved extremities for measurement of cytokines interleukin (IL)-6, IL-18 and tumor necrosis factor- $\alpha$  (TNF- $\alpha$ ), the neuropetides NPY and CRGP, and prostaglandin E2. Results: The patients included in this study did have a moderate to serious disease activity and impairment. In plasma, no changes of mediators of inflammation were observed In blister fluid, however, significantly higher levels of IL-6 and  $TNF-\alpha$  in the involved extremity were observed in comparison with the uninvolved extremity. Conclusions: This is the first time that involvement of mediators of inflammation in CRPS 1 has been so clearly and directly demonstrated. This observation opens new approaches for the successful use and development of immunosuppressives in CRPS 1. 2002:305303 HCAPLUS <<LOGINID::20081016>> AN 137:167971 DN ΤI Evidence for local inflammation in complex regional pain syndrome type 1 AU Huygen, Frank J. P. M.; De Bruijn, Anke G. J.; De Bruin, Martha T.; Groeneweg, J. George; Klein, Jan; Zijlstra, Freek J. CS Pain Treatment Centre, Erasmus Medical Centre, Rotterdam, 3000 CA, Neth. SO Mediators of Inflammation (2002), 11(1), 47-51 CODEN: MNFLEF: ISSN: 0962-9351 PB Taylor & Francis Ltd. DT Journal LA English RE.CNT 32 THERE ARE 32 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 4 OF 4 HCAPLUS COPYRIGHT 2008 ACS on STN

TI Increased production of nitric oxide stimulated by interferon-γ from peripheral blood monocytes in patients with complex regional pain syndrome

AB This study examines immediate nitric oxide (NO) release from monocytes following interleukin-1β (IL-1β), interferon-γ (IFN-γ), and tumor necrosis factor-.

alpha. (TNF- $\alpha$  ) challenge in patients

with complex regional pain syndrome

(CRPS). Study patients exhibited the following: (1), mech. allodynia; (2), evidence of either vasomotor or sudomotor disturbance; and (3), concordant painful allodynia documented with quant. sensory testing that was temporarily abolished with sympathetic block. Ten subjects (CRPS,

N=5; control, N=5) were enrolled. Peripheral blood monocytes were challenged with 100  $\mu L$  of IL-IB (I ng), IFN-y (1 ng), TFN-c (0.01 ng), and normal saline (NS) and the resultant immediate NO release measured. Subjects with CRPS exhibited a statistically significant increase in NO release in response to IFN-y compared with controls. The NO responses to IFN-y in excess of NS and as the ratio IFN-y/NS were also significantly increased.

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- DN 136:368210
- TI Increased production of nitric oxide stimulated by interferon-γ from peripheral blood monocytes in patients with complex regional pain syndrome
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- CS Department of Anesthesiology and Perioperative Medicine, William Beaumont Hospital, Royal Oak, MI, 48073, USA
- SO Neuroscience Letters (2002), 323(1), 75-77 CODEN: NELED5; ISSN: 0304-3940
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